DSSTox Log File:

FDA Maximum (Recommended) Daily Dose Database (FDAMDD)

(last updated 10 April 2006)

Description: Information in this file documents creation, review, and update process for the DBPCAN SDF file, provides summary information on database content, and lists currently unavailable CAS registry numbers for known structures. The first section summarizes the process used for creating the initial DSSTox SDF files and the quality assurance checks and procedures employed. A table providing field and data counts offers summary overview of DBPCAN file content. The Log Table documents modifications and revisions to the database content or format in version updates. To obtain the most current version of this Log File and a record of any new modifications, or to report errors in this file, a user should consult the DSSTox DBPCAN database page: http://www.epa.gov/nheerl/dsstox/sdf dbpcan.html

Description: Information in this file documents the creation, review, and update process for the DSSTox FDAMDD SDF files and provides summary information on database contents. The first section summarizes the process used for creating the initial DSSTox SDF file and the quality assurance checks and procedures employed. A table providing field and data counts offers summary overview of FDAMDD file contents and chemical composition. The Log Table documents modifications and revisions to the database content or format in version updates. To obtain the most current version of this Log File and a record of any new modifications, a user should periodically consult the DSSTox FDAMDD database page: http://www.epa.gov/nheerl/dsstox/sdf fdamdd.html.

QA and Development Notes for v1a:

FDAMDD SDF file underwent an extensive series of quality review checks prior to publication of initial launch version. The original FDA MRDD database was obtained from the Source website (http://www.fda.gov/cder/Offices/OPS IO/MRTD.htm), imported into Excel, and was used for cross-referencing and checking of all modified and added fields. We particularly thank Edwin Matthews, Daniel Benz, and Naomi Kruhlak for invaluable assistance in various stages of development and quality review, clarifying numerous issues pertaining to the experimental data and their representation in the original database, and approving changes to field names and contents. Chemical structures were initially obtained by conversion of the FDA Source-provided SMILES codes using CambridgeSoft ChemDraw (ver 8.0 for Windows) for MS Excel. An additional field containing CAS registry numbers was contributed to this effort by the FDA Source collaborators. However, CAS were not provided for all listed chemicals and, in many cases, multiple CAS were provided for single structure records in the original Source database. For the latter, the CAS corresponding to the main SMILES structure was often not apparent since additional CAS often corresponded to salt and complexed forms or different stereoisomer forms of the listed pharmaceutical. Hence, considerable additional effort was invested in the verification, review, and addition of CAS registry numbers to the DSSTox FDAMDD database.

Four sources of CAS registry numbers (CASRN) corresponding to chemical structures were used extensively in the checking of chemical structures and CASRN, identification of missing CASRNs, and identification of derivative forms of structures corresponding to alternate CASRN listed within a structure entry. The ChemFinder website (http://chemfinder.cambridgesoft.com/) and National Library of Medicine's ChemID website (<a href="http://chem.sis.nlm.nih.gov/chemidplus/chemidplu

IUPAC systematic chemical names, **ChemName_IUPAC**, were computed by Marc Nicklaus (NCI) using the ACD Labs IUPAC Name-Generation software (ACD/NameBatch, version 8.05). **INChI** codes were automatically generated from the final DSSTox SDF using a pre-release version of the publicly available program, wINChI11b.exe, accessible from the NIST INChI developers (http://chemdata.nist.gov/IChI/INChIv11b.zip).

A total of 15 records were deleted in migrating the original Source-provided FDA MRDD database (1233 records as of 20May04) to the DSSTox FDAMDD (1217 final records in version 1a). These 15 records are listed in a table below along with the original FDA Source SMILES, the problem with the record, and the action taken in creating the initial launch version of the DSSTox FDAMDD.

All records in the DSSTox FDAMDD SDF file are "defined organics" (i.e., no inorganics or organometallics). A file in which all salts and complexes are simplified to the parent structure can be generated easily using commercial and publicly available tools from the information provided in the DSSTox FDAMDD SDF file.

Notes for v2a,b:

Revised DSSTox Standard Chemical Fields are included (see http://www.epa.gov/nheerl/dsstox/MoreonStandardChemFields.html) along with updated InChI codes (version 1.0), recomputed IUPAC chemical names (ACDLabs ACD/Name, version 8.0), and many regenerated 2D structures with stereochemistry of steroidal compounds rendered in more standardized form. Additionally, an extensive quality review of all DSSTox chemical records was performed, resulting in numerous corrections and modifications to chemical structures and added information (CASRN, representative structures for mixtures, etc) throughout DSSTox data files. For more information on current review procedures, see http://www.epa.gov/nheerl/dsstox/ChemicalInfQAProcedures.html

Log of SDF Modifications and Version/revision updates:

Date	DSSTox SDF File Names	Modifications from previous version	Additional Notes
1Mar05	FDAMDD_v1a_1217_1Mar05	Initial launch publication; no previous published versions.	FDAMDD is considered a "static" historical database, although further alterations or expansion of the database is possible. Future updates will primarily correct reported errors provided by users or incorporate DSSTox format changes.
10Apr2006	FDAMDD_v2b_1217_10Apr2006	Updated with new DSSTox Standard Chemical Fields and entries (revised Aug 2005). Updated InChI codes (version 1.0). Updated IUPAC chemical names (ACDLabs Name to Structure, version 8.0). Expanded "ddmmmyear" format for dates in DSSTox file names (e.g., 15Dec2005). Modified Source-related field names: TherapeuticCategory ChemClass_MRDD_grouping ActivityCategory_MRDD_mmol ActivityCategory_MCASE_mg Deleted field SMILES_Parent_nostereo	Numerous structure modifications and changes in stereochemical rendering throughout DSSTox data files following major quality review. FDAMDD_v2a_1217_22Oct2005: Note: earlier version of this file was provided to PubChem, with identical format to v2b but latter has undergone additional QA review and has small number of corrections/modifications.

Field and Data Counts in the DSSTox FDAMDD SDF file:

DSSTox SDF	Standard Standard Chemical Toxicity Fields Fields		Source- specific fields	Chemical records total	STRUCTURE_ ChemicalType:	STRUCTURE_TestedForm_ DefinedOrganic:		
					Defined organic	Parent	Salt	Complex
FDAMDD_v1	20	3	8	1217*	1217	1137	58	22

^{*} One pair of replicate parent structures (Bephenium), and two pairs of replicate CAS (Iophendylate, m and o and Iodinated Glycerol A and B) exist in the database.

DSSTox SDF	TestSubsatnce _CASRN	TestSubstance_ CASRN_Other	TestSubstance_ CASRN_Other (identified in ChemicalNote)	ChemicalNote (stereochem)	Chemical Note (related forms) 2	ChemClass_ MRDD_ grouping (group classes) 3	Chemical Replicate Count=1 ⁴
FDAMDD_v1	1217	1277	911	483	107	37	1147

⁴ Number of "unique" structure/activity entries in FDAMDD, where only the first member of an **ChemClass_MRDD_grouping** is counted.

FDAMDD SDF Content*	Totals_v2b	
# Records	1217	
DSSTox Standard Chemical Fields	20	
DSSTox Standard Toxicity Fields	3	
FDAMDD Source Fields	8	
Total # Fields	31	

¹ Not including duplicate listings of CAS for related forms.

² Entry "related forms" indicates record is member of an **ChemClass_MRDD_grouping**, sharing a common MRDD value with a number of closely related structural derivatives.

³ Number of groups where a general drug family name is assigned to closely related structural derivatives sharing a common **Dose_MRDD_mg** value.

Chemical Content	Counts_v2b
STRUCTURE_ChemicalType:	
defined organic	1217
inorganic	0
organometallic	0
no structure	0
STRUCTURE_TestedForm_ DefinedOrganic:	
parent	1136
complex	24
salt	7
salt complex	0
TestSubstance_Description:	
single chemical compound	0
defined mixture or formulation	1216
undefined mixture	0
macromolecule	1
unspecified or multiple forms	0

Records Deleted from FDA Source MRDD Database (ver. 20May04) to create DSSTox FDAMDD_v1a:

Source Chemical Name	Source SMILES	Reason for deletion of record	Action
Acetyldigitoxin	C1C(O)CC2C(C)(C1)C3C(CC2)C5(O)C(C)(CC3)C(C4=CC(=O)OC4)CC5, C1(C)C(O)C(O)CC(O)O1	Original Source database listed two records with this ChemName, each containing only a portion of the full structure SMILES.	List full structure in single record of FDAMDD.
Carbinoxamine	C2=CC=C(C(OCCN(C)C)C1=CC=C(CI)C=C1)N=C2, S(=O)(=O)(OS(=O)(=O)OC1=CC=CC=C1)OC2=CC=CC=C2	Original Source database listed two records with this ChemName, the first Carbinoxamine and the second resembling besylate. No CAS or Merck entry located for any form resembling source SMILES complex; CAS provided by source was for maleate complex.	List parent form in FDAMDD with correct CAS.
Cisatracurium (2 records deleted)	C1(OC)=C(OC)C=C6C(=C1)CC[N+](C)(CCC(=O)OCCCCCO C(=O)CC[N+]3(C)C(CC2=CC=C(OC)C(OC)=C2)C4=C(CC3)C =C(OC)C(OC)=C4)C6CC5=CC(OC)=C(OC)C=C5, C1=CC=CC=C1S(O)(=O)=O	Original Source database listed two records with this ChemName, the first Cisatracurium and the second a partial representation of besylate. Atracurium is same chemical as Cisatracurium, but incorrect structure SMILES was listed in original Source database (extra CH3).	Delete both Cisatracurium records and replace Atracurium parent with besylate complex form in FDAMDD.
Deanol aceglumate	C(C)(=O)NC(CCC(O)=O)C(=O)O	Source indicated that Dose_MRDD_mg dose value corresponds exclusively to Deanol listing.	Delete record.
Dichloralphenazone	C(CI)(CI)(CI)C(O)O, C2=CC=CC(N1C(=O)C=C(C)N1C)=C2	Original Source database listed two records with this ChemName, the first entry corresponding to single chloral entity, the second to phenazone portion.	List as single record in complex form with 2 chlorals.
Difluoromethylornithine, alpha-	NCCCC(N)(C(O)O)C(F)F	Structure appears incorrect and no CAS provided by Source; name is synonym for Eflornithine, which is listed as separate record.	Delete record as duplicate of Eflornithine.
Diminazenediaceturate	OC(=O)CNC(C)=O, C2(C(=N)N)=CC=C(NN=NC1=CC=C(C(N)=N)C=C1)C=C2	Original Source database listed two records with this ChemName, the first the diaceturate moiety and the second the parent diminazene.	List as single record in complex form.
Homatropine	C3=CC=CC(C(O)C(=O)OC1CC2N(C)C(C1)CC2)=C3	Source indicated that Dose_MRDD_mg dose value corresponds exclusively to Homatropine, dl methylbromide listing.	Delete record.
Mebrobamate	NC(=O)OCC(C)(CCC)COC(N)=O	Name appears incorrect and no CAS provided by Source. Also, Source SMILES only slightly different from SMILES for Meprobamate record; latter is presumed correct.	Delete record for Mebrobamate.
Methyclothiazide	C1(S(=O)(=O)N)=C(C)C=C2C(=C1)S(O)(O)N(C)C(CCI)N2	Structure from Source SMILES appears incorrect; correct structure but incorrect	Delete record for Methyclothiazide and

		name listed for Metychlothiazide record.	correct name in Metychlothiazide record (to Methyclothiazide).
Phenylsemicarbazide, 1-	NC(NNC1=CC=CC=C1)=O	Original Source database listed wrong structure for this name; structure the same as for Phenicarbazide.	Delete record.
Pyruvinium Pamoate	C3(C(O)=O)=CC4C(C(CC1C(O)=C(C(=O)O)C=C2C=1C=CC=C2)=C3O)=CC=CC=4	Original Source database listed two records with this ChemName, the first the neutral Pamoate moiety and the second the parent Pyruvinium.	List as single record in complex form.
Sulbactam, t-butanoic acid	C1C(=O)N2C1S(=O)(=O)C(C)(C)C2C(=O)OCOC(=O)C(C)(C) C	Source indicated that Dose_MRDD_mg dose value corresponds exclusively to Sulbactam listing.	Delete record.
Thioquanine	N1C(N)=NC2=C(C1=S)NC=N2	Name appears incorrect and no CAS provided by Source. Structure is tautomeric form of Thioguanine listed as separate record.	Delete record for Thioquanine.
Tribromoethanol, 1,2,2	C(Br)(Br)C(O)Br	Source indicated that Dose_MRDD_mg dose value corresponds exclusively to Tribromoethanol, 2,2,2 listing.	Delete record.